The function of phosphatidylinositol 5-phosphate 4-kinase γ (PI5P4Kγ) explored using a specific inhibitor that targets the PI5P binding site.

Jonathan H. Clarke*†, Maria-Luisa Giudici*†, John E. Burke‡§, Roger L. Williams‡, David J. Maloney¶, Juan Marugan¶ & Robin F. Irvine*

* Department of Pharmacology, Tennis Court Road, Cambridge CB2 1PD, UK
‡MRC Laboratory of Molecular Biology, Francis Crick Avenue, Cambridge CB2 0QH, UK
¶National Center for Advancing Translational Sciences, 9800 Medical Center Drive, Rockville, MD 20850, USA
§ Present address: Department of Biochemistry and Microbiology, University of Victoria, 270a Petch Hall, Victoria, BC, Canada
† These authors contributed equally to this study.

Page heading title: Phosphatidylinositol 5-phosphate 4-kinase γ inhibition

Corresponding author, R.F. Irvine rfi20@cam.ac.uk  Phone 44-1223-334177 FAX 44-1223 334040

Keywords: Phosphatidylinositol 5-phosphate; phosphatidylinositol 5-phosphate 4-kinase γ; PI5P; PI5P4K; PI5P4Kγ

Abbreviations: PI5P, phosphatidylinositol 5-phosphate; PI(4,5)P2, phosphatidylinositol 4,5-bisphosphate; HDX-MS, Hydrogen-deuterium exchange mass spectrometry.
Abstract

NIH-12848 (NCGC00012848-02), a putative phosphatidylinositol-5-phosphate 4-kinase γ (PI5P4Kγ) inhibitor, was explored as a tool for investigating this enigmatic, low activity, lipid kinase. PI5P 4-kinase assays \textit{in vitro} showed that NIH-12848 inhibited PI5P4Kγ with an IC$_{50}$ of approximately 1µM but did not inhibit the α and β PI5P4K isoforms at concentrations up to 100µM. A lack of inhibition of PI5P4Kγ ATPase activity suggested that NIH-12848 does not interact with the enzyme’s ATP binding site, and direct exploration of binding using hydrogen-deuterium exchange mass spectrometry revealed the putative PI5P-binding site of PI5P4Kγ to be the likely region of interaction. This was confirmed by a series of mutation experiments, which led to the identification of a single PI5P4Kγ amino acid residue that can be mutated to its PI5P4Ks α and β homologue to render PI5P4Kγ resistant to NIH-12848 inhibition. 10µM NIH-12848 was applied to cultured mouse principal kidney cortical collecting duct (mpkCCD) cells, which we show express PI5P4Kγ that increases when the cells grow to confluence and polarise. NIH-12848 inhibited the translocation of Na$^+$/K$^+$ATPase to the plasma membrane that occurs when mpkCCD cells grow to confluence, and also prevented reversibly their forming of ‘domes’ on the culture dish. Both these NIH-12848-induced effects were mimicked by specific RNAi knock-down of PI5P4Kγ, but not that of PI5P4Ks α or β. Overall the data reveal a likely contribution of PI5P4Kγ to the development and maintenance of epithelial cell functional polarity, and show that NIH-12848 is a potentially powerful tool for exploring the cell physiology of PI5P4Ks.
Introduction

The phosphatidylinositol 5-phosphate 4-kinases (PI5P4Ks) are a family consisting of three isoforms in mammals (α, β, and γ), whose most likely function is currently believed to be the regulation of the levels of their substrate, PI5P (see refs [1, 2] for reviews). While the α and β isoforms have had some physiological and pathological functions assigned to them, including in gene regulation [3, 4], stress responses [5], insulin signalling [6] and cancer [7, 8], PI5P4Kγ is hardly understood at all. It is apparently ubiquitously expressed, but at very different levels between tissues, being particularly high in epithelial cells of the thick ascending limb of the kidney and collecting ducts cells [9] and in specific neurons in brain [10]. It apparently has a vesicular localization [9, 10], may have a functional connection with Rho [11], shows a much lower enzymatic activity in vitro than the other PI5P4K isoforms [12], and can heterodimerise in vitro with PI5P4Kα [12] (note that extensive heterodimerisation between PI5P4Ks α and β has been shown to occur in vivo [13, 14]).

Specific inhibitors of enzymes can be useful tools in studying their function, and kinase inhibitors are among those that have shown most promise as potential therapeutic agents. Recently the characterization of inhibitors for PI5P4Kα [15] and PI5P4Kβ [16] have raised that hope for those isoforms, but the isoform specificity of neither inhibitor has yet been established, and so far no such tools exist at all for PI5P4Kγ. Moreover, a challenge facing any kinase inhibitor, the great majority of which interact with the ATP-binding site of their target, is for it to have both sufficient specificity (because all kinase ATP binding sites show some structural similarity) and potency (cellular concentrations of ATP are in the millimolar range, so nanomolar affinity of an inhibitor is often required for micromolar efficacy in a cell). The high affinity and specificity of the PI 3-kinase δ inhibitor PIK-39 that results from a remarkable induced fit into the ATP binding site of its target protein [17] is one example of an ATP binding site competitor that overcomes these issues. A potential approach for increasing the kinase specificity is to look for ATP-allosteric modulators, although in some cases (e.g. [18]) there are discrepancies between cell-based and isolated kinase inhibitory assays, making difficult the finding of this kind of inhibitor.

Herein we report the characterization and use of a PI5P4Kγ-specific inhibitor NIH-12848 (full designation NCGC00012848-02), which we show interacts not with the ATP binding site but with the region where PI5P probably binds, including the activation loop. We use the inhibitor to begin the first exploration, in a kidney epithelial cell line, of the function of PI5P4Kγ. Also, we show how we can mutate PI5P4Kγ so that it becomes insensitive to NIH-12848, opening the possibility of chemical biology to explore the functions of all three PI5P4Ks.

Materials and Methods

Enzyme preparation and mutagenesis

Recombinant enzyme was prepared essentially as described previously [12]. Protein from PIP4K2C (UniGene 6280511) or associated mutants, cloned into the expression vector pGEX6P (GE Healthcare) was expressed and purified from E. coli
BL21(DE3). Cultures were induced with 0.4 mM IPTG and probe-sonicated in the presence of protease inhibitors. GST fusion proteins of PI5P4Kγ and PI5P4Kγ+, a mutant with specific activity close to that of the active PI5P4Kα isoform [12], were harvested by binding to glutathione sepharose beads (GE Healthcare) and cleaved in situ with 50U of PreScission protease (GE Healthcare) for 4 hours at 4°C. Purity was confirmed by SDS-PAGE and protein concentration determined by colorimetric assay (Bio-Rad). Site-directed mutagenesis using the QuikChange technique (Agilent Technologies) was used to generate clones from which mutant forms of PI5P4Kγ and PI5P4Kγ+ were produced (for mutagenesis primers see Table S1).

Biochemical assays

Lipid kinase assays were performed as described previously [13]. DiC16-PI5P was purchased from Echelon Biosciences, and after drying down in Eppendorf tubes was sonicated for 3 x 30 seconds in a Decon Ultrasonics sonicating bath. This lipid substrate (6 µM PI5P) and recombinant lipid kinase were added to the reaction mixture (200µl final volume) with 10 µCi [γ-32P]ATP and incubated at 30°C for 10-60 min. Lipids were extracted using an acidic phase-separation [19] and separated by one-dimensional thin layer chromatography (2.8:4:1:0.6 chloroform:methanol:water:ammonia). Radiolabelled PI(4,5)P2 spots were detected by autoradiography, extracted and counted with Ultima Gold XR scintillant (Packard) on a LS6500 scintillation counter (Beckman Coulter). Intrinsic ATPase activities of the enzymes were determined using the Transcreener ADP² fluorescence polarization method (BellBrook Labs). A range of enzyme concentrations was assayed with ATP substrate (100µM ATP, 60 min incubation at 22°C) and polarization units (mP) were read using a PHERAstar Plus microplate reader (BMG Labtech). Experimental values were interpolated from an ADP/ATP utilization standard curve and plotted using nonlinear regression analysis with Prism 5.

Hydrogen Deuterium Exchange Mass Spectrometry

Hydrogen deuterium exchange mass spectrometry experiments were performed according to previously published protocols [20, 21]. In brief, HDX reactions were initiated in HDX buffer (20 mM HEPES pH 7.5, 50 mM NaCl, 2 mM Tris Carboxyl Ethyl Phosphine (TCEP)) by the addition of 98% D2O Solution (10 mM HEPES pH 7.5, 50 mM NaCl, 2 mM TCEP), to give a final concentration of 78% D2O. Final protein concentrations were 1 µM. For all reactions with inhibitor there was a ten fold excess of inhibitor to protein, and this was allowed to incubate for 30 minutes on ice before initiation of H/D exchange. Four time points of exchange were carried out (3 s, 30 s, and 300 s at 23 °C and 3 s at 0 °C), and were terminated by the addition of a quench buffer (1 M guanidine-HCl, 0.8% formic acid), followed by snap freezing in liquid nitrogen. Samples were stored at -80 °C until mass analysis.

Measurement of Deuterium Incorporation

Protein samples were rapidly thawed and injected onto a UPLC column immersed in ice as previously described [21]. The protein was run over two immobilized pepsin columns in series (Applied Biosystems; porosyme, 2-3131-00) at 200 µL/min for 3 min and collected over a van-guard pre-column trap (Waters). The trap was subsequently eluted in line with an Acquity 1.7 µm particle, 100 mm x 1 mm C18 UPLC column (Waters), using a gradient of 5-36% B (buffer A 0.1% formic acid, buffer B 100% acetonitrile) over 20 min. Eluent from the column was injected
onto a Xevo QTOF (Waters) acquiring over a mass range from 350 to 1500 m/z, with an electrospray ionization (ESI) source operated at a temperature of 225 °C, and a spray voltage of 2.5 kV.

Protein Digestion, Peptide Identification, and Mass Analysis

Peptide identification was done by running tandem MS/MS experiments using a 3-35% B gradient over 120 min with a Xevo QTOF (Waters). This was supplemented with a 20 min gradient separation to identify and correct the retention time for all samples. The MS tolerance was set to 3 ppm with an MS/MS tolerance at 0.1 Da. The resulting MS/MS datasets were analyzed with the Mascot search within Mascot distiller (matrix science). All peptides with a Mascot score >15 were analyzed using HD-Examiner Software (Sierra Analytics). The full list of peptides was then manually validated by searching a non-deuterated protein sample’s MS scan to test for the correct m/z state, and check for the presence of overlapping peptides. Ambiguously identified peptides were excluded from all subsequent analysis. The first round of analysis and identification were performed automatically by the HD-Examiner software, but all peptides (deuterated and non-deuterated) were manually verified at every state and time point for the correct charge state, m/z range, presence of overlapping peptides, and the expected retention time. All HDX-MS results are presented as relative levels of deuterium incorporation, and no correction for back exchange is applied, because no fully deuterated protein sample could be obtained.

Cell Culture

Immortalized mouse cortical collecting duct (mpkCCD) cells [22, 23] were grown in defined medium (DMEM:Ham’s F12 1:1 vol/vol, 60 nM sodium selenate, 5 µg/ml transferrin, 2 mM glutamine, 50 nM dexamethasone, 1 nM triiodothyronine, 10 ng/ml epidermal growth factor, 5 µg/ml insulin, 20 mM D-glucose, 2% vol/vol fetal calf serum, and 20 mM HEPES, pH 7.4) at 37°C in a 5% CO₂-95% air atmosphere. Medium was changed every two days and all studies were performed on cells that had been grown on plastic Petri dishes or glass slides.

RNAi

Depletion of specific PI5P4K isoforms was achieved by transfecting cells with small interference RNAs (siRNAs) specific to each PI5P4K isoform. The siRNAs were obtained from Thermo Scientific Dharmacon (ON-TARGET plus SMART pool). Control siRNAs were sequences that would not be recognised by any part of the mouse genome. MpkCCD cells were plated at a density of 6x10⁴ per well (24 well/plate) one day before transfection, then were transfected with each siRNA at a final concentration of 25 pmol/ml using DharmaFECT 1 Transfection Reagent (Dharmacon). At 72-h post-transfection, a new round of transfection was performed, and the cells were cultured for another 48 h.

Morphological and and immunocytochemical studies

Confluent cells grown on plastic Petri dishes were examined under an inverted microscope (Zeiss) equipped with phase-contrast optics. Indirect immunofluorescence studies were performed on confluent cells grown on glass slides fixed with ice-cold methanol or 4% formaldehyde. Cells were processed for immunofluorescence using anti-Na⁺/K⁺-ATPase α-1 monoclonal antibody (Millipore) and anti-ZO-1 polyclonal antibodies (Millipore), and were examined under a Leica confocal microscope.
Western Blotting

Cells were grown to confluence, washed in cold PBS, and lysed by scraping them off in boiling Laemmli sample buffer containing 20mM β-mercaptoethanol. Lysate was boiled for 5 min, and viscosity was reduced by sonication using four bursts of 30 sec each with 15 sec on ice between each sonication step. Lysate was boiled as before and centrifuged at 20000xg for 5 min. The supernatant was run on 10% poly-acrylamide gels, transferred to nitrocellulose membranes, blocked in TBS-0.05% Tween 20/5% non-fat dry milk and probed with antibodies diluted in the same solution. The following primary antibodies were used: polyclonal anti-PI5P4Kγ 1:2500 [9], monoclonal anti-α tubulin 1:5000 (SIGMA). After washing in TBS-0.05% Twen and incubation with secondary antibody, protein was detected by Super Signal West Femto Maximum Sensitivity Substrate (Thermo Scientific).

QT-PCR

Cellular RNA was extracted with Trizol reagent (Invitrogen) and 1 ng was reverse transcribed with Superscript III and random hexamers (Invitrogen). Quantification of gene expression of the three PI5P4K isoforms was assessed by using TaqMan Gene Expression Assays and TaqMan Gene Expression Master Mix (Life Technologies), using TATA-box binding protein (TBP) as the reference gene.

Results and Discussion

NIH-12848 is a specific PI5P4Kγ inhibitor

NIH-12848 (Fig 1) initially emerged as ‘compound 1’ from a kinome profiling study, measuring its selectivity against a panel of 460 kinases (see ref [24] Supplementary Figure 3). As can be seen in the supporting information (Supplementary Table S2), NIH-12848 was exquisitely selective for PI5P4Kγ, with an apparent IC_{50} of 2-3μM (Fig S1). This screen did not include PI5P4Kα (Fig S2), and to clarify the actions of NIH-12848 against PI5P4Ks we tested it in PI5P 4-kinase assays using PI5P and 32P-γ-ATP as substrates (see Methods). Under these conditions, PI5P4Kα was not inhibited by 100μM NIH-12848 (Fig S3A), and PI5P4Kβ showed a small but significant stimulation at the same concentration (Fig S3B), whereas PI5P4Kγ was inhibited with an apparent IC_{50} of 3.3μM (Fig 2), similar to the apparent Kd in the kinome profiling (above).

Because PI5P4Kγ has a very low catalytic activity [12] these assays use a final enzyme concentration of around 2.3 μM, so it is possible that NIH-12848 is actually more potent and we are in part titrating the enzyme protein. We therefore also tested NIH-12848 on nanomolar levels of the PI5P4Kγ+ mutant [12], which has a 300-fold higher activity than wild type PI5P4Kγ, and we found the IC_{50} of NIH-12848 to be about 1μM (Supp Fig 3C). Below we show that the PI5P binding site is the likely site of interaction for NIH-12848, and because the lipid PI5P does not have a precise concentration (not least this depends on its physicochemical structure), the number we deduce here can only be an empirical IC_{50}, not an absolute K_{i}.

NIH-12848 interacts with the PI5P binding site on PI5P4Kγ

Most of the mutations introduced into PI5P4Kγ to create PI5P4Kγ+ [12] involve the changing of residues that bind or stabilise ATP to the equivalent residues present in the much more active PI5P4Kα. So the marked contrast in sensitivities to NIH-12848 of PI5P4Kγ+ versus PI5P4Kα suggested that NIH-12848 could be
unusual among kinase inhibitors in that it might not be interacting with the ATP
binding site. This possibility was supported by an inability of NIH-12848 to inhibit
the intrinsic ATPase activity of PI5P4Kγ (Fig 2B), and by the observation that
including 100µM ATP in assays using PI5P4Kγ+ (Km for ATP, 2µM [12]) had only a
small effect on the ability of NIH-12848 to inhibit the enzyme near its IC50
concentration: 2µM NIH-124848 inhibited PI5P4Kγ+ by 40±3.2% with carrier-free
(approx. 16 nM) ATP, and 64±2.8% with 100µM ATP; the small shift in NIH-12848
sensitivity is probably an allosteric effect (see below).

To determine directly the interaction site of NIH-12848 with PI5P4Kγ and
PI5P4Kγ+ we turned to hydrogen deuterium exchange mass spectrometry (HDX-
MS), a powerful tool for mapping protein conformational changes, and binding
epitope sites of interaction on proteins [21, 25]. There are two peptides in PI5P4Kγ
whose rate of HDX is altered significantly in the presence of NIH-12848 (Fig 3B-D).
One peptide spans residues 373-407 (Fig 3C,D), part of the enzyme’s activation loop,
which has been shown to govern the substrate specificity of PI5P4Ks and PI4P5Ks
[26, 27]. There was also a change in a peptide spanning residues 158-162 (Fig 3C,D),
which is part of an α-helix that is near the activation loop in PI5P4Kγ [28], designated
α-helix 4 in the original annotated PI5P4Kβ structure ([29]), see Fig 3B,F).

Intriguingly, in the PI5P4Kγ+ there was a significant increase in exchange in residues
158-162 compared to wild-type PI5P4Kγ, and a correspondingly larger decrease in
exchange upon addition of NIH-12848 (Fig 3E). These data point to the proposed
PI5P-binding site of PI5P4Ks [26, 27, 29] as the likely area of NIH-12848 interaction.

**PI5P4Kγ can be made resistant to NIH-12848**

Direct competition experiments (increasing PI5P and looking at the effect that
this has on NIH-12848 potency *in vitro*) would be the most direct route to confirming
the above suggestion. However, NIH-12848 is a hydrophobic compound, as is PI5P,
and it is likely that the two compounds would form mixed micellar structures when
together in aqueous buffers, so any such ‘competition’ would be uninterpretable. To
seek confirmation for the site of interaction of NIH-12848 we therefore undertook a
series of mutational experiments, focusing on the regions of interaction suggested by
the HDX-MS data. In the region 373-407 (see above), only glutamine 378 differs
between PI4P5K isoforms and mutating this residue to the PI5P4Kγ equivalent
histidine made no detectable difference to the activity or NIH-12848 sensitivity of the
enzyme (data not shown). In residues 158-162 (see above), isoleucine 159 and
aspartate 161 are both valines in PI5P4Ks α and β, and mutating these together in
PI5P4Kγ+ caused a small shift in sensitivity to NIH-12848 (inhibition by 10µM NIH-
12848 was 96%±1.5 in WT PI5P4Kγ+ and 71%±2.6 in the (E378H,I159V,D161V)
PI5P4Kγ+ mutant).

Examining this region closely in the structure (Fig 3F) suggested three other
residues that might interact with the substrate or inhibitor and which are different
between PI5P4Ks γ versus α/β. These are serine 164, asparagine 165, and glutamate
153, which are respectively asparagine, isoleucine and threonine in PI5P4Kα/β; of
these, glutamate 153 has already been mutated to the PI5P4Kα/β-equivalent threonine
in creating PI5P4Kγ+ [12]. We therefore introduced all five mutations of potential
interest (Q378H, I159V, D161V, S164N, N165I) into PI5P4Kγ and PI5P4Kγ+, and
both these mutant enzymes were completely resistant to NIH-12848 inhibition (Fig
S4A,B).

These data support the HDXMS data suggesting that NIH-12848 is binding to
this area of the PI5P4Kγ protein, and they also imply that serine 164 and/or
asparagine 165 are the major contributors to the sensitivity of PI5P4Kγ to NIH-12848. So we mutated only those two residues in PI5P4Kγ to their PI5P4Ks α/β homologues individually or together, and found that while S164N PI5P4Kγ is as sensitive to NIH-12848 inhibition as wild type PI5P4Kγ (Fig S4C), N165I PI5P4Kγ is completely insensitive, with 50µM NIH-12848 having no effect on this mutant (Fig 4). Note that the activity of N165I PI5P4Kγ was 6.7x higher than wild type PI5P4Kγ in these assays, and Fig S4D summarises the activities and sensitivities to NIH12848 of the mutants discussed here.

Overall these data show that asparagine 165 is the residue in PI5P4Kγ that is almost entirely responsible for its sensitivity to NIH-12848. There was no detectable change in the amide exchange rate at N165 when NIH-12848 binds to PI5P4Kγ or PI5P4Kγ+ (Fig 3), and the simplest interpretation is that NIH-12848 binds to the activation loop and α-helix 4 to induce a change in the structure in this region, and that introducing an isoleucine (as in PI5P4Ks α and β) at residue 165 prevents this structural change from happening, perhaps because of a hydrophobic interaction with the activation loop (see Fig 3F). Whatever the molecular mechanism, this residue could in the future be mutated in both alleles of the PI5P4Kγ gene in a cell line or a mouse to engender NIH-12848 resistance of the PI5P4Kγ, which would be a powerful chemical biology approach to eliminate completely any potential non-specific cellular effects of NIH-12848. Introducing the complementary mutations into PI5P4Ks α or β, possibly to make them NIH-12848-sensitive, could be an equally informative avenue of investigation.

Effects of NIH-12848 on a kidney cell line

PI5P4Kγ is widely expressed in mammalian tissues, but is particularly high, and is indeed the principal PI5P4K isoform, in kidney [9]. It is expressed there in epithelial cells, particularly in the thick ascending limb and cortical collecting duct, so to explore its cellular functions we chose mpkCCD cells, a cell line derived from epithelial cells in the cortical collecting duct of the kidney [22, 23]. Western blotting confirmed an easily detectable level of expression of PI5P4Kγ in mpkCCD cells (Fig 6A). These cells grow to confluence and then form gap junctions between them. They then begin to form ‘domes’ in which process, they show a property of transporting epithelia when cultured in vitro on solid supports. ‘Domes’ are fluid-filled blisters formed between the solid growth surface and the cell layer, and their formation is regarded as a sign of differentiated epithelia with active transport processes and with an intact epithelial barrier due to functional cell-cell contacts [22, 23]. We found using quantitative PCR that while reaching confluence there is a 3-fold increase in PI5P4Kγ mRNA, but no significant change in the two other PI5P4K isoforms (Table 1). We thought that PI5P4Kγ might be involved in epithelial cell polarity because in adult mammalian kidney PI5P4Kγ is present in intracellular vesicles with a distribution polarised towards the lumen [9], and in zebra fish (Danio) it is highly expressed in the pro-nephric duct throughout development (ZFIN Database: ZDB-IMAGE-030829-506).

One easily monitored parameter is the redistribution of Na+/K+ ATPase from the cytoplasm to the basolateral plasma membrane that occurs when the cells reach confluence [30]. We explored the effect of 10µM NIH-12848 (1µM had similar but smaller effects) and found a complete inhibition of this redistribution, while the cells remained healthy and confluent, with gap junctions formed as revealed by immunocytochemistry of ZO-1 (Fig 5A). From this, we suggest PI5P4Kγ is involved in the progress towards a polarised epithelial phenotype.
Consistent with this possibility, the formation of ‘domes’ (see above) was also markedly inhibited (Fig 5B) by NIH-12848. Moreover, if NIH-12848 was added subsequent to dome formation, the domes disappeared with a detectable effect within 3 hours that was maximal by 24 hours (Fig 5B), indicating a dynamic relationship between PI5P4Kγ and the development and maintenance of this morphology. The molecular mechanisms underlying this relationship will require much further investigation, but the reported link between PI5P4Kγ and Rho [11], taken together with the well-known involvement of Rho in epithelial cell transport and polarity [31, 32], offers one possibility for exploration.

**RNAi knock-down of PI5P4Kγ**

Our extensive molecular exploration of the interaction of NIH-12848 with PI5P4Kγ (above) emphasizes its remarkable specificity, but nevertheless it is possible that the effects we see are due to an off-target action. To gather some independent evidence that inhibition of PI5P4Kγ underlies the effects of NIH-12848, and thus to begin a validation of this inhibitor as a tool for exploring PI5P4Kγ function, we knocked down PI5P4Kγ in the mpkCCD cells by RNAi. Fig 6A and Table S3 shows that the knock-down was specific and successful, and Figs 6 B-C show the effects of PI5P4Kγ knock-down on Na+/K+ATPase re-localisation (Fig 6B) and dome formation (Fig 6C). Within the limits of these analyses, both effects of NIH-12848 are mimicked by this knock-down.

Control experiments knocking down PI5P4Kα and β did not show the same effect on dome formation. PI5P4Kβ knock-down had no detectable effect (Fig S5B), while knockdown of PI5P4Kα had a pronounced opposite (stimulatory) effect (Fig S5A,B). We have no immediate explanation for this latter action, which we believe is likely to be independent of PI5P4Kγ activity because knock-down of PI5P4Kγ and PI5P4Kα together showed that PI5P4Kγ causes a similar and internally consistent inhibitory effect on the higher level of dome formation resulting from PI5P4Kα knock-down (Fig S5C). Further studies, including using PI5P4Kα inhibitors (e.g. ref [15]) will be required to clarify this function of PI5P4Kα and its relationship with PI5P4Kγ. We have discussed elsewhere [12] the possibility that PI5P4Kγ may sometimes serve to target the more active PI5P4Kα to a specific cellular location.

Overall these experiments yield the first functional insight into PI5P4Kγ’s physiological role in epithelial cells. Much remains to be explored to clarify exactly what the enzyme is doing in this context, but the very similar effects induced in mpkCCD cells by inhibition with NIH-12848 and knock-down of PI5P4Kγ are a crucial first stage in a proof of principle of the potential usefulness of NIH-12848 in such an exploration. Our detailed mapping and manipulation of the molecular interaction of NIH-12848 with PI5P4Kγ also suggests ways to a wider use of this inhibitor in the exploration of functions of the PI5P4K family.

**Acknowledgements**

We are very grateful to Dr A. Vandewalle (INSERM U773, Paris, France) for provision of the mpkCCD cell line and Simon Bulley for his help and advice with the QT-PCR. J.H.C was supported by the MRC (Grant RG64071), M-L.G. by the BBSRC (Grant RG65394), and J.H.B. by the BHF (Grant PG11/109/29247).
Author contributions:

Jon Clarke engineered and produced recombinant proteins, performed in vitro assays and processed data; Maria-Luisa Giudici performed cell biology experiments and QT-PCR and processed data; John Burke performed the HDX-MS, processed data and interpreted protein structures; Roger Williams processed HDX-MS data and interpreted protein structures; David Maloney and Juan Marugan performed the initial discovery of the PI5P4Kγ-NIH-12848 interaction; Robin Irvine performed some in vitro assays and processed data. All the authors discussed the results, interpreted data and co-wrote the paper.

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Figure Legends

Figure 1 Structure of NIH-12848

Figure 2 Effect of inhibitor NIH-12848 on PI5P4Kγ activity. A) Dose-response curve showing inhibition of PI5P 4-kinase activity over a range of inhibitor concentrations (0.01 to 100 µM). Points represent a mean of 5 replicates and error bars indicate ± S.E.M. B) Effect of a range of inhibitor concentrations on intrinsic ATPase activity of PI5P4Kγ. Bars represent % ATP conversion by PI5P4Kγ (1µM) in the presence of NIH-12848, in comparison to the indicated positive control (●; 1µM PI5P4Kγ in the absence of inhibitor). A negative control was also included (□; inhibitor only, no enzyme). N=3, error bars represent ± S.E.M.

Figure 3 Binding of NIH-12848 to PI5P4Kγ studied by HDX-MS. A) Global H/D exchange of Apo PI5P4Kγ. B) Structure of PI5P4Kγ with changes induced by 10µM NIH-12848 shown in pseudo-colour. C) Summary of changes induced by NIH-12848. D) Kinetics of H/D exchange ± 10µM NIH-12848 in two regions of PI5P4Kγ. E) Quantification of H/D exchange in region of residues 158-162 in PI5P4Kγ and PI5P4Kγ+ with or without NIH-12848. F) Model of PI5P4Kγ active site [28] with residues mutated in this study highlighted in red (note that the E153T mutation engineered in PI5P4Kγ+ [12] is highlighted here in yellow – see text). The activation loop (containing Gln378) is not visible in the X-ray structure, and is a conjectural conformation, as is the location of the ATP and PI5P substrates. Also in yellow (but not indicated) are Asp374, which is a conserved aspartate crucial for ATP binding [29], and Ala386, which dictates the substrate specificity of PI5P4Ks [26, 27].

Figure 4 Effect of inhibitor NIH-12848 on PI5P4Kγ harbouring a single amino acid substitution. PI5P4Kγ mutant N165I was assayed in the presence of 0, 5 and 50 µM NIH-12848 and showed no significant sensitivity towards this inhibitor. N= 3-6 replicates for each condition, error bars represent S.E.M.

Figure 5 Effects of NIH-12848 on mpkCCD cells. A) Effect of 10µM NIH-12848 on Na⁺/K⁺ ATPase α−1 localization. Cells were grown on glass coverslips at sub-confluence and then treated for 24 h with inhibitor or carrier (DMSO). They were stained for ZO-1 (gap junctions) or Na⁺/K⁺ ATPase α-1 and viewed under a confocal microscope. Low resolution imaging showed that >90% of the cells were as shown in these higher resolution images, and at least three experiments were performed with identical results. Objective 100X. Bars, 10µm. B) Effect of 10µM NIH-12848 on dome formation. Cells were grown on plastic dishes to confluence (no domes present) and then for a further 48 hours with inhibitor or carrier. Phase contrast images of cell monolayers are shown. Objective 20X. Bars, 50μm. C) Effect of 10µM NIH-12848 after domes have formed. Cells were grown on plastic dishes for 5 days to allow dome formation and then with inhibitor or carrier for a further 24 hours. Phase contrast images of cell monolayers are shown. Objective 20X. Bars, 50 µm.
Figure 6

A) Specificity and depletion efficiency of PI5P4Kγ siRNA. mpkCCD cells grown on plastic dishes were transfected with control siRNA or PI5P4Kγ-specific siRNA. After 72 hours, the transfection was repeated. After further 48 hours, RNA was extracted and the expression levels of each PI5P4K isoform was evaluated by RT-qPCR. Values (above) are expressed as fold difference in knocked-down cells relative to control cells with TATA-box binding protein (TBP) as the reference gene. Depletion of PI5P4Kγ was also evaluated in cell lysates by Western blot (below). Equal volumes of cell lysates were loaded on a 10% SDS-PAGE gel and PI5P4Kγ was detected with an isoform-specific polyclonal antibody (see Methods).

B) Effect of siRNA knock-down of PI5P4Kγ on Na+/K+-ATPase α−1 localization. Cells grown on glass coverslips were depleted of endogenous PI5P4Kγ by siRNA as described in Material and Methods. Confocal sections of cells fixed and stained for Na+/K+-ATPase α−1 (green) and ZO-1 (red) are shown. These images are typical of two independent experiments, and the images are representative of >90% of cells examined. Objective 63X. Bars, 10 µm.

C) Effect of siRNA knock down of PI5P4Kγ on dome formation. mpkCCD cells grown on plastic dishes were depleted of endogenous PI5P4Kγ by siRNA as described in Material and Methods. Phase contrast images of cell monolayers are shown. Objective 10X. Bars, 50 µm.

D) Surface density of cell domes was measured in mpkCCD confluent cell monolayers that were transfected with PI5P4Kγ-specific siRNA or only transfection reagent (Control 2) or Non Target Oligos siRNA (Control 3). In Control 1 cells were not transfected. Values are expressed as means ± SEM. *p<0.001 versus Control 3. Significant differences were analyzed by Student’s t-test.
Table 1

mRNA levels of PI5P4Ks in mpkCCD cells grown to confluence

<table>
<thead>
<tr>
<th>Gene</th>
<th>PI5P4Kγ</th>
<th>PI5P4Kα</th>
<th>PI5P4Kβ</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polarized mpkCCD cells</td>
<td>3.19-3.46</td>
<td>0.99-1.01</td>
<td>0.51-0.53</td>
</tr>
</tbody>
</table>

The relative expression of the three genes PI5P4Kγ, α, and β, with TATA-box binding protein as a reference gene, was determined in a RT-qPCR assay. Values are expressed as fold difference in polarized cells (four days after reaching confluence when domes are formed) relative to pre-confluent cells (no domes present).
Figure 1

NIH-12848
Figure 2

A

specific activity (nmol ATP converted into PI5P min⁻¹ mg⁻¹)

\log [\text{inhibitor}], \mu M

B

% conversion of ATP

[\text{inhibitor}], \mu M
Figure 3

HDX-MS

Fig 3A

Fig 3B

Fig 3C

Fig 3D
Figure 4

![Graph showing specific activity versus μM inhibitor](image-url)
Figure 5

Fig 5A

Na⁺/K⁺ ATPase α−1  
ZO-1

Control

+ NIH-12848
Figure 6

Fig 6A

<table>
<thead>
<tr>
<th>Gene</th>
<th>PI5P4Kγ</th>
<th>PI5P4Kα</th>
<th>PI5P4Kβ</th>
</tr>
</thead>
<tbody>
<tr>
<td>Relative mRNA expression</td>
<td>0.16-0.18</td>
<td>0.92-0.98</td>
<td>0.81-0.84</td>
</tr>
</tbody>
</table>

siRNA control  siRNA PI5P4Kγ

PI5P4K

Tubulin
Fig 6B
Na+/K+-ATPase α-1    ZO-1

siRNA control

siRNA-PI5P4Kγ
**Supplementary Tables**

**Supplementary Table 1.**
Primer pairs that were used to generate amino acid substitutions in PI5P4Kγ and PI5P4Kγ+ mutants.

<table>
<thead>
<tr>
<th>Amino acid change</th>
<th>Primer pair (forward/reverse)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Q378H</td>
<td>5'-TATCCTTACACATTATGATGCCAAGAAG</td>
</tr>
<tr>
<td>I159V + D161E</td>
<td>5'-CCAGTGAGGACGTTGCTGAAATGCATAGCAAC</td>
</tr>
<tr>
<td>S164N</td>
<td>5'-TGACATGCATAACACCTCTCAAACATG</td>
</tr>
<tr>
<td>N165I</td>
<td>5'-TGACATGCATAGCATCCTCTCAAACATCAC</td>
</tr>
</tbody>
</table>

**Supplementary Table 2**

Data from Kinome screen (see xls file)

**Supplementary Table 3**

mRNA levels of PI5P4Ks following RNAi induced knock down.

<table>
<thead>
<tr>
<th>Relative mRNA expression</th>
<th>PI5P4Kγ</th>
<th>PI5P4Kα</th>
<th>PI5P4Kβ</th>
</tr>
</thead>
<tbody>
<tr>
<td>si-RNA-PI5P4Kγ</td>
<td>0.16-0.18</td>
<td>0.92-0.98</td>
<td>0.81-0.84</td>
</tr>
<tr>
<td>siRNA-PI5P4Kα</td>
<td>1.10-1.40</td>
<td>0.15-0.20</td>
<td>1.00-1.10</td>
</tr>
<tr>
<td>siRNA-PI5P4Kβ</td>
<td>1.00-1.20</td>
<td>0.84-0.98</td>
<td>0.30-0.32</td>
</tr>
</tbody>
</table>

mpkCCD cells grown on plastic dishes were transfected with control siRNA or PI5P4K–specific siRNAs. After 72 hours, transfection was repeated. After further 48 hours, RNA was extracted and the expression levels of each PI5P4K isoform was evaluated by RT-qPCR.
Supplementary Figures

Supplementary Figure Legends

Figure S1
Dose-response of NIH-12848 on PI5P4Kγ in Kinome screen.

Figure S2
Data for PIP kinases extracted from Supplementary Table S2

Figure S3
Effect of inhibitor NIH-12848 on PI5P4K activity. Specific activities of PI5P4Kα (A) and PI5P4Kβ (B) are not inhibited when assayed in the presence of NIH-12848. N=3, error bars represent ±S.E.M. C) Dose response curve of PI5P4Kγ+ inhibited by NIH-12848. Values represent mean reduced activity compared to the mean of uninhibited PI5P4Kγ+ replicates from the same experiment.

Figure S4
Effect of inhibitor NIH-12848 on the activity of PI5P4Kγ mutants. A) Specific activity of PI5P4Kγ and B) PI5P4Kγ+, both with five amino acid substitutions (Q378I, I159V, D161E, S164N and N165I) were assayed in the presence of 0, 5 and 50 µM NIH-12848 inhibitor. C) Specific activity of PI5P4Kγ with a single amino acid substitution (S164N) in the presence of 1, 2 and 5 µM NIH-12848 inhibitor. D) Summary of the specific activities of a full range of PI5P4Kγ mutants in the presence of no (-) or 5 µM (+) NIH-12848. Substituted amino acids are listed below the relevant bars. WT = wild type PI5P4Kγ. N= a minimum of 3 replicates for each experiment, error bars represent S.E.M.

Figure S5
Effects of RNAi knockdown of PI5P4Kα and β on dome formation in mpkCCD cells. A) siRNA for PI5P4Kα or control were applied in two rounds over 5 days. Typical images are shown. B) Quantification of experiments typified by Fig S5A (with data for equivalent experiments with siRNA for PI5P4Kβ) Values are expressed as means ± SEM. *p<0.001. Significant differences were analyzed by Student’s t-test. C) Quantification of experiments in which PI5P4Kα and γ were knocked down alone or together. Values are expressed as means ± SEM. *p<0.001. Significant differences were analyzed by Student’s t-test.
Supplementary Figures

Figure S1

Figure S2
Figure S3

(A) Specific activity (error bars indicate SD)

(B) Specific activity (error bars indicate SD)

(C) % reduced activity

log[Inhibitor], μM
Figure S4
Figure S5

S5A
siRNA-control  siRNAi-PI5P4Kα

S5B

![Bar chart showing the number of domes/mm² for different groups](chart.png)

- Control
- siRNA-PI5P4Kα
- siRNA-PI5P4Kβ

* indicates a significant difference.
Fig S5C

![Bar graph showing Nr domes/mm² for different conditions: Control, siRNA-PSP4Ag, siRNA-PSP4Ag/β, and siRNA-PSP4Ag/canty. The graph includes error bars and asterisks (*) indicating statistical significance.](image)